

April 13, 2015

The Honorable Fred Upton, Chairman  
House Committee on Energy & Commerce  
2125 Rayburn House Office Building  
Washington, D.C. 20515

The Honorable Frank Pallone, Ranking Member  
House Committee on Energy & Commerce  
2322A Rayburn House Office Building  
Washington, D.C. 20515

Dear Chairman Upton and Ranking Member Pallone:

On behalf of the 30 million men, women and children in the U.S. living with a rare disease, the undersigned organizations urge you to permanently authorize the *Rare Pediatric Disease Priority Review Voucher (PRV) Program* to drive greater development of novel treatments for children with a rare pediatric disease.

There are an estimated 7,000 rare diseases, which are defined as a disease affecting 200,000 or fewer people. Of the nearly one in ten Americans with a rare disease, approximately two-thirds are children. Of the 350 most "common" rare diseases, 27 percent result in death before the child's first birthday.

Despite significant unmet medical need (the approximately 450 approved orphan products treat only about 350 rare diseases), manufacturers face significant obstacles that can hinder the pursuit of rare disease therapies for children, including difficulties associated with conducting clinical trials. To tackle these hurdles, Congress established the *Rare Pediatric Disease PRV Program*.

Currently, upon FDA approval of a novel rare pediatric disease treatment, the *Rare Pediatric Disease PRV Program* provides a biopharmaceutical manufacturer the opportunity to receive a voucher guaranteeing a six month priority review of a New Drug Application (NDA) or Biologic License Application (BLA) for another product, rare disease or not. The voucher can be sold to another company, and there is no limit on how often it may be transferred.

Unfortunately, this program expires March 2016, ending a clear pathway that encourages innovators to pursue treatments in a difficult disease space. To date, three vouchers have been awarded and the program has shown clear evidence that it is a valuable incentive to develop drugs and biologics in this underserved area.

Congress established the *Rare Pediatric Disease PRV Program* because it recognized the necessity of an incentive to enhance innovation in this key area of unmet patient need, a market segment previously overlooked.

We urge Congress to permanently authorize the *Rare Pediatric Disease PRV Program*, which has proven its initial effectiveness in providing hope to children who are suffering from these rare conditions, and drawing manufacturers to invest in the development of novel treatments for rare pediatric diseases.

Sincerely,

Adult Polyglucosan Body Disease Research Foundation  
 ALD Connect  
 Alstrom Angels  
 Alstrom Syndrome International  
 American Association of the Deaf-Blind  
 American Autoimmune Related Diseases Association  
 American Partnership For Eosinophilic Disorders  
 American Thoracic Society  
 Amyloidosis Support Groups Inc  
 Association for the Bladder Exstrophy Community  
 Association for Creatine Deficiencies  
 Association for Glycogen Storage Disease  
 Autoinflammatory Alliance  
 Avery's Angels Gastroschisis Foundation  
 Batten Disease Support and Research Association  
 Barth Syndrome Foundation  
 Bridge the Gap  
 CADASIL Together We Have Hope Non-Profit Organization  
 Canavan Foundation  
 CARES Foundation  
 CCHS Family Network  
 CFC International  
 Charcot-Marie-Tooth Association  
 Children's Brittle Bone Foundation  
 Children's Cardiomyopathy Foundation  
 Children's PKU Network  
 Chronic Granulomatous Disease Association  
 Circadian Sleep Disorders Network  
 Coalition for Pulmonary Fibrosis  
 Congenital Hyperinsulinism International (CHI)  
 Cooley's Anemia Foundation  
 Council for Bile Acid Deficiency Diseases  
 CureCADASIL Association  
 Cure AHC  
 Cure HHT  
 Cure JM Foundation  
 Cure SMA  
 debra of America  
 The Dent Diseases Foundation  
 Dravet Foundation  
 Dupuytren Foundation  
 EB Research Partnership  
 Encephalitis Global  
 Everylife Foundation for Rare Diseases  
 Fabry Support & Information Group  
 Fight ALD-Fighting Illness Through Education

FOD (Fatty Oxidation Disorders) Family Support Group  
 Foundation Fighting Blindness  
 Foundation for Angelman Syndrome Therapeutics  
 Foundation for Ichthyosis & Related Skin Types  
 FPIES Foundation  
 Friedreich's Ataxia Research Alliance  
 Galactosemia Foundation  
 GBS/CIDP Foundation International  
 Global Genes  
 Gwendolyn Strong Foundation  
 Hereditary Neuropathy Foundation  
 Histiocytosis Association  
 Hope for Hypothalamic Hamartomas  
 International FOP Association  
 International Pemphigus and Pemphigoid Foundation (IPPF)  
 Jeffrey Modell Foundation  
 LGS Foundation  
 Lipodystrophy United  
 Little Miss Hannah Foundation  
 Lymphangiomatosis & Gorham's Disease Alliance  
 Lymphedema Advocacy Group  
 The Marfan Foundation  
 Moebius Syndrome Foundation  
 Myotonic Dystrophy Foundation  
 National Adrenal Diseases Foundation  
 National Alopecia Areata Foundation  
 National Ataxia Foundation  
 National Brain Tumor Society  
 National Eosinophilia Myalgia Syndrome Network  
 National MPS Society  
 National Organization for Rare Disorders  
 National Stem Cell Foundation  
 National Tay-Sachs & Allied Diseases Association  
 NBIA Disorders Association  
 NGLY1.org  
 Noah's Hope Fund  
 NTM Info & Research  
 Organic Acidemia Association  
 Oxalosis and Hyperoxaluria Foundation  
 Parents and Researchers Interested in Smith-Magenis Syndrome  
 PCDH19 Alliances  
 PF Strategies  
 Phelan-McDermid Syndrome Foundation  
 Pituitary Network Association  
 Potocki-Lupski Syndrome Outreach Foundation  
 Project DOCC - Delivery of Chronic Care

Pulmonary Hypertension Association  
Rare and Undiagnosed Network (RUN)  
Rare Disease United Foundation  
Rare Genomics Institute  
The Reflex Sympathetic Dystrophy Syndrome Association (RSDSA)  
Rettsyndrome.org  
Sanfilippo Foundation for Children  
Scleroderma Foundation  
Simons VIP Connect  
Stickler Involved People  
Sturge-Weber Foundation  
Tarlov Cyst Disease Foundation  
The Transverse Myelitis Association  
The United Leukodystrophy Foundation  
United Mitochondrial Disease Foundation  
U.S. Hereditary Angioedema Association  
Usher Syndrome Coalition  
Vascular Birthmarks Foundation  
VHL Alliance  
We are RARE Inc.  
XLH Network  
The XLP Research Trust  
5p-Society

For additional information, contact Paul Melmeyer, Assistant Director of Public Policy, National Organization for Rare Disorders (NORD), [pmelmeyer@rarediseases.org](mailto:pmelmeyer@rarediseases.org), (202) 588-5700 ext. 104.